

Sequences producing significant alignments:	Score (bits)	E Value	
gi 23238206 ref NM_014452.3 Homo sapiens tumor necrosis fa	417		GIU,E
gi 37181727 gb AY358304.1 Homo sapiens clone DNA52594 DR6	<u>417</u>	e-114	
gi 7022799 dbj AK001504.1 Homo sapiens cDNA FLJ10642 fis, gi 6478163 emb AL096801.18 HSJ181J13 Human DNA sequence fro gi 15880636 emb AJ336218.1 HSA336218 Homo sapiens genomic s gi 15880407 emb AJ335989.1 HSA335989 Homo sapiens genomic s	417 417 409 325	e-114 e-111 3e-86	GWE
gi 15868604 emb AJ324225.1 HSA324225 Homo sapiens genomic s	<u>269</u>	2e-69	abre
gi 34783298 gb BC017730.2 Homo sapiens tumor necrosis fact	<u>166</u>		G U E
$\frac{\text{gi} 3549262 \text{gb} \text{AF068868.1} }{\text{gi} 32880108 \text{gb} \text{BT009883.1} } \text{Homo sapiens TNFR-related death r}$	$\frac{144}{144}$	1e-31	GÜE
$\begin{array}{ll} \underline{\text{gi} 30583678 \text{gb} \text{BT007420.1} } & \text{Homo sapiens tumor necrosis fact} \\ \underline{\text{gi} 55626753 \text{ref} \text{XM}} & \underline{518523.1} & \text{PREDICTED: Pan troglodytes tum} \\ \underline{\text{gi} 15870715 \text{emb} \text{AJ326321.1} \text{HSA326321}} & \text{Homo sapiens genomic s} \\ \end{array}$	$\frac{144}{136}$ $\frac{122}{122}$	1e-31 3e-29 5e-25	G Ü]E
gi 34874517 ref XM 236992.2 Rattus norvegicus similar to d	<u>96</u>	4e-17	GU
gi 31341673 ref NM 178589.2 Mus musculus tumor necrosis fa	88	9e-15	GU
<pre>gi 26335926 dbj AK043823.1 Mus musculus 10 days neonate co</pre>	88	9e-15	
<pre>gi 16741136 gb BC016420.1 Mus musculus tumor necrosis fact</pre>	86	4e-14	G UE
$\begin{array}{lll} \underline{\text{gi} 26329206 \text{dbj} \text{AK033529.1} } & \text{Mus musculus adult male colon c} \\ \underline{\text{gi} 15877202 \text{emb} \text{AJ332784.1} \text{HSA332784}} & \text{Homo sapiens genomic s} \\ \underline{\text{gi} 17432071 \text{emb} \text{AL646086.1} } & \text{Ralstonia solanacearum GMI1000} & \dots \end{array}$	82 52 44	6e-13 6e-04 0.15	GlU

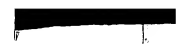
		11	
<pre>gi 56178122 gb AE017340.1 Idiomarina loihiensis L2TR, comp gi 22795236 gb AC127460.2 Homo sapiens chromosome 5 clone gi 14625084 gb AC084380.8 AC084380 Oryza sativa chromosome</pre>	42 42 42	0.59 0.59 0.59	
gi 33879154 gb BC021165.2 Homo sapiens zinc finger protein	40	2.3	GU
gi 33870187 gb BC013011.2 Homo sapiens zinc finger protein	40	2.3	GU
gi 34871182 ref XM 343874.1 Rattus norvegicus similar to f	40	2.3	GU
gi 25453488 gb AY057439.2 Azotobacter vinelandii GroEL (gr gi 52421840 gb CP000011.1 Burkholderia mallei ATCC 23344 c	40	2.3	
<pre>gi 34222201 ref NM 032772.3 Homo sapiens zinc finger prote gi 52211453 emb BX571966.1 Burkholderia pseudomallei strai</pre>	40	2.3	GU
<u>gi 34534651 dbj AK127647.1 </u> Homo sapiens cDNA FLJ45745 fis,	40	2.3	GU
<u>gi 21751602 dbj AK092897.1 </u> Homo sapiens cDNA FLJ35578 fis,	40	2.3	GU
gi 21106326 gb AE011651.1 Xanthomonas axonopodis pv. citri	40	2.3	G OU'E
<pre>gi 33871437 gb BC011625.2 Homo sapiens zinc finger protein gi 20303531 gb AC010997.12 Homo sapiens chromosome 10 clon</pre>	40	2.3	GUE
<u>gi 15875552 emb AJ331134.1 HSA331134</u> Homo sapiens genomic s gi 18653743 gb AC012236.12 Homo sapiens chromosome 15, clo	$\frac{40}{40}$	2.3	
gi 29608854 dbj AP005042.1 Streptomyces avermitilis genomi	40	2.3	
gi 24646031 ref NM 141841.1 Drosophila melanogaster CG1053 gi 23171004 gb AE003692.3 Drosophila melanogaster chromoso	38	9.4 9.4	GU
gi 31208238 ref XM_313086.1 Anopheles gambiae ENSANGP00000	_38	9.4	GU
gi 50947384 ref XM 483220.1 Oryza sativa (japonica cultiva	_38	9.4	G
<pre>gi 51965001 ref XM_507285.1 PREDICTED Oryza sativa (japoni</pre>	_38	9.4	U
<pre>gi 50929414 ref XM_474235.1 Oryza sativa (japonica cultiva</pre>	_38	9.4	G
<pre>gi 34896883 ref NM_184897.1 Oryza sativa (japonica cultiva</pre>	<u>38</u>	9.4	G
gi 37534057 ref NM_196349.1 Oryza sativa (japonica cultiva	_38	9.4	G
<u>gi 50937328 ref XM 478192.1 </u> Oryza sativa (japonica cultiva <u>gi 28269488 gb AC092781.6 </u> Oryza sativa chromosome 3 BAC OS	<u>38</u> <u>38</u>	9.4 9.4	G
<u>gi 29789306 ref NM 057200.1 </u> Rattus norvegicus kinesin fami	38	9.4	GU
<u>gi 31431860 gb AE017090.1 </u> Oryza sativa (japonica cultivar <u>gi 32489031 emb AL606999.3 OSJN00127</u> Oryza sativa genomic D	<u>38</u> <u>38</u>	9.4 9.4	
gi 42469651 emb BX820650.1 CNSOAA3D Arabidopsis thaliana Fu	_38	9.4	U
gi 42468881 emb BX818953.1 CNS0A9UM Arabidopsis thaliana Fu	_38	9.4	U
gi 42468460 emb BX820086.1 CNS0A9DI Arabidopsis thaliana Fu	_38	9.4	U
gi 42468388 emb BX819814.1 CNS0A9FM Arabidopsis thaliana Fu	_38	9.4	
gi 42468355 emb BX819698.1 CNSOA9GB Arabidopsis thaliana Fu gi 22296994 gb AC105101.8 Homo sapiens chromosome 18, clon	<u>38</u> 38	9.4 9.4	U
gi 38257164 gb AC109601.9 Oryza sativa chromosome 3 BAC OS	38	9.4	
<pre>gi 40786619 gb AC096856.7 Oryza sativa chromosome 3 BAC OS gi 42408124 dbj AP004190.3 Oryza sativa (japonica cultivar</pre>	<u>38</u> 38	9.4	
<u>gi 21428993 gb AY119562.1 </u> Drosophila melanogaster GM13228	38	9.4	Ü
gi 13421727 gb AE005727.1 Caulobacter crescentus CB15 sect	38	9.4	G
gi 21387210 gb AY114690.1 Arabidopsis thaliana proline imi	38	9.4	GW
gi 37988923 dbj AK119300.1 Oryza sativa (japonica cultivar	38	9.4	U.
<pre>gi 21111436 gb AE012143.1 Xanthomonas campestris pv. campe gi 37806382 dbj AP006064.3 Oryza sativa (japonica cultivar</pre>	<u>38</u> <u>38</u>	9.4 9.4	G
gi 9947599 gb AE004591.1 Pseudomonas aeruginosa PAO1, sect	38	9.4	G
gi 21689936 emb AL390778.31 Human DNA sequence from clone	38	9.4	
gi 32984072 dbj AK098863.1 Oryza sativa (japonica cultivar	38	9.4	M M
<pre>gi 32982206 dbj AK072183.1 Oryza sativa (japonica cultivar</pre>	<u> 38</u>	9.4	U

]
I
1
l
1
ŧΕ
•
ı

Alignments

Cetselected sequences And Selectell Deselectell

Sbjct: 299 agtgcacatggggtgttggaggtagatgggctcccgggcccgggaggcggcggtggatgcg 358



```
Query: 133 gcgctgggcag-mgcagccgccgattccagctgccccgcgcgcccc-ggccaccttgcga 190
        Sbjct: 359 gcgctgggcagaagcagccgccgattccagctgccccgcgcgcccccgggcgcccctgcga 418
Query: 191 gtccccggttcagccatggggacctctccgagcagcagcaccgccctcggcctcctgcm- 249
        Sbjct: 419 gtccccggttcagccatggggacctctccgagcagcagcaccgccctc-gcctcctgcag 477
Query: 250 ccqcatcqccqqqqccacaqccacqatqatcqcqqqctccct 294
        Sbjct: 478 ccgcatcgccqcqaqccacaqccacqatqatcqcqqqctccct 522
Length = 3534
Score = 417 bits (209), Expect = e-114
Identities = 270/285 (94%), Gaps = 6/285 (2%)
Strand = Plus / Plus
Query: 15 gcmggtgctgagcgcccctagngcctcccttgccgcctccctctctctgccggccgtagc 74
        Sbjct: 130 gcaggtgctgagcgcccctagagcctcccttqccqcctcctctctctqccqqccqcaqc 189
Query: 75 aqtgcacat-gngtgttggaggtagatgggctcccgg-ccgggaggcggcggtggatgcg 132
        Sbjct: 190 agtgcacatggggtgttggaggtagatgggctcccggcccgggaggcggcggtggatgcg 249
Query: 133 gcgctgggcag-mgcagccgccgattccagctgccccgcgcgcccc-ggccaccttgcga 190
        Sbjct: 250 gcgctgggcagaagcagccgccgattccagctgccccgcgcgcccccgggcgcccctgcga 309
Query: 191 gtccccggttcagccatggggacctctccgagcagcagcaccgccctcggcctcctgcm- 249
        Sbjct: 310 gtccccggttcagccatggggacctctccgagcagcagcaccgccctc-gcctcctgcag 368
Query: 250 ccgcatcgcccgccgagccacagccacgatgatcgcgggctccct 294
        Sbjct: 369 ccgcatcgcccgccgagccacagccacgatgatcgcgggctccct 413
- >qi|7022799|dbj|AK001504.1| GUE Homo sapiens cDNA FLJ10642 fis, clone NT2RP200
        to Homo sapiens TNFR-related death receptor-6 mRNA
       Length = 2636
Score = 417 \text{ bits } (209), \text{ Expect} = e-114
Identities = 270/285 (94%), Gaps = 6/285 (2%)
Strand = Plus / Plus
```

```
Query: 15 gcmggtgctgagcgcccctagngcctcccttgccgcctccctctctcccggccgtagc 74
       Query: 75 agtgcacat-gngtgttggaggtagatgggctcccgg-ccgggaggcggcggtggatgcg 132
       Sbjct: 260 agtgcacatggggtgttggaggtagatgggctcccgggcccgggaggcggcggtggatgcg 319
Query: 133 gcgctgggcag-mgcagccgccgattccagctgccccgcgcgcccc-ggccaccttgcga 190
       Sbjct: 320 gcgctgggcagaagcagccgctgttccagctgccccgcgcgcccccgggcgcccctgcga 379
Query: 191 gtccccggttcagccatggggacctctccgagcagcagcaccgccctcggcctcctgcm- 249
       Sbjct: 380 gtccccggttcagccatggggacctctccgagcagcagcaccgccctc-gcctcctgcag 438
Query: 250 ccgcatcgccgcgagccacagccacgatgatcgcgggctccct 294
       Sbjct: 439 ccgcatcgcccgccgagccacagccacgatgatcgcgggctccct 483
Contains the TNFRSF21 gene for tumor necrosis factor
        receptor superfamily superfamily member 21 (DR6) and a CpG
        Island, complete sequence
      Length = 118271
Score = 417 \text{ bits } (209), \text{ Expect = } e-114
Identities = 270/285 (94%), Gaps = 6/285 (2%)
Strand = Plus / Plus
Query: 15
        gemggtgetgagegeeectagngeeteeettgeegeeteeeteetetgeeeggeegtage 74
        {\tt agtgcacat-gngtgttggaggtagatgggctcccgg-ccgggaggcggtggatgcg} \ \ 132
Query: 75
         Sbjct: 36822 agtgcacatggggtgttggaggtagatgggctcccgggcccgggaggcggcggtggatgcg 36881
Query: 133
        gcgctgggcag-mgcagccgccgattccagctgccccgcgcgcccc-ggccaccttgcga 190
        Sbjct: 36882 gcgctgggcagaagcagccgccgattccagctgccccgcgcgcccccgggcgcccctgcga 36941
Query: 191
        gtccccggttcagccatggggacctctccgagcagcagcaccgccctcggcctcctgcm- 249
        Sbjct: 36942 gtccccggttcagccatggggacctctccgagcagcagcaccgccctc-gcctcctgcag 37000
Query: 250
        ccgcatcgccgccgagccacagccacgatgatcgcgggctccct 294
```

```
Sbjct: 37001 ccgcatcgcccgaccacaccaccacgatgatcgcgggctccct 37045
☐ >gi|15880636|emb|AJ336218.1|HSA336218
                                Homo sapiens genomic sequence surrounding
        NL6-FL22C
       Length = 666
 Score = 409 bits (205), Expect = e-111
 Identities = 269/285 (94%), Gaps = 6/285 (2%)
 Strand = Plus / Plus
Query: 15 gcmggtgctgagcgccctagngcctcccttgccgcctcctctctctgccggccgtagc 74
        Sbjct: 17
        gcaggtgctgagcgccctagagcctcccttqccqcctcctctctctqcccqccqcaqc 76
Query: 75 agtgcacat-gngtgttggaggtagatgggctcccgg-ccgggaggcggtggatgcg 132
        Sbjct: 77 agtgcacatggggtgttggaggtagatgggctcccgggcccgggaggcggccgatgcg 136
Query: 133 gcgctgggcag-mgcagccgccgattccagctgccccqcgcgcccc-qqccaccttqcqa 190
        Sbjct: 137 gegetgggeagaageageegeegatteeagetgeeeegegegeeeegggegeeetgega 196
Query: 191 gtccccggttcagccatggggacctctccgagcagcagcaccgccctcggcctcctgcm- 249
        Sbjct: 197 gtccccggttcagccatggggacctctccgagcagcagcaccgccctc-gcctcctqcaq 255
Query: 250 ccgcatcgcccgccgagccacagccacgatgatcgcgggctccct 294
        Sbjct: 256 ccgcatcgcccgccgagccacagccacgatgatcgcgggctccct 300
. Sqi|15880407|emb|AJ335989.1|HSA335989
                                Homo sapiens genomic sequence surrounding
        NL6-DN19C
       Length = 693
Score = 325 \text{ bits (163)}, Expect = 3e-86
Identities = 257/283 (90%), Gaps = 6/283 (2%)
Strand = Plus / Plus
Query: 15 gcmggtgctgagcgccctagngcctcccttgccgcctcctctctctgccggccgtagc 74
        Sbjct: 17
        Query: 75 agtgcacat-gngtgttggaggtagatgggctcccgg-ccgggaggcggtggatgcg 132
        Sbjct: 77 agtgcacatggggtgttggaggtagatgggctcccgggcccgggaggcggccgatgcg 136
```

```
Query: 133 gcgctgggcag-mgcagccgccgattccagctgccccgcgcgcccc-ggccaccttgcga 190
        Sbjct: 137 gcgctgggcagaaacagccgccgattccaactgccccgcgcgccccgggcgcccctgcga 196
Query: 191 gtccccggttcagccatggggacctctccgagcagcagcagcaccgcctcggctcctgc-m 249
        Sbjct: 197 gtccccggttcagccatggggacctttccgagcagcagcacctgcctc-gccttctgcaa 255
Query: 250 ccgcatcgccgccgagccacagccacgatgatcgcgggctcc 292
        Sbjct: 256 ccgcatcgaccgccgagccacattcacgatgatcgctggctcc 298
☐ >gi|15868604|emb|AJ324225.1|HSA324225
                                Homo sapiens genomic sequence surrounding
        NL1-ZP16C
       Length = 762
Score = 269 \text{ bits } (135), \text{ Expect} = 2e-69
Identities = 202/221 (91%), Gaps = 4/221 (1%)
Strand = Plus / Plus
Query: 15 gcmggtgctgagcgcccctagngcctcccttgccgcctccctctctctgccggccgtagc 74
        Query: 75 agtgcacat-gngtgttggaggtagatgggctcccgg-ccgggaggcggcqgtggatgcg 132
        Sbjct: 77 agtgcacatggggtgttggaggtagatgggctcccggcccgggaggcggtcgatgcg 136
Query: 133 gcgctgggcag-mgcagccgccgattccagctgccccgcgcgcccc-ggccaccttgcga 190
        Sbjct: 137 gcgctgggcagaagcagccgccgattccagctgccccgcgcgccccggggcgcccctgcga 196
Query: 191 gtccccggttcagccatggggacctctccgagcagcagcac 231
        Sbjct: 197 gtcctgggctcagccatggggacctctgagaggagcagcac 237
□>qi|34783298|qb|BC017730.2| GUE Homo sapiens tumor necrosis factor receptor su
        mRNA (cDNA clone MGC:21476 IMAGE:3847246), complete cds
       Length = 3238
Score = 166 \text{ bits } (83), \text{ Expect = } 3e-38
Identities = 98/101 (97%), Gaps = 2/101 (1%)
Strand = Plus / Plus
Query: 195 ccggttcagccatggggacctctccgagcagcagcaccgccctcggcctcctgcm-ccgc 253
        ccggttcagccatggggacctctccgagcagcagcaccgccctcq-cctcctqcaqccqc 59
```

```
Query: 254 atcgcccqccqaqccacagccacqatgatcqcgggctccct 294
         Sbjct: 60 atcgcccgccgagccacagccacgatgatcgcgggctccct 100
                         GUE Homo sapiens TNFR-related death receptor-6 (DR6
\square >qi|3549262|qb|AF068868.1|
        Length = 1968
 Score = 144 bits (72), Expect = 1e-31
 Identities = 87/90 (96%), Gaps = 2/90 (2%)
 Strand = Plus / Plus
Query: 206 atggggacctctccgagcagcagcaccgccctcggcctcctgcm-ccgcatcgcccgccg 264
         Sbict: 1
         atggggacctctccgagcagcagcaccgccctcg-cctcctgcagccgcatcgccgccg 59
Query: 265 agccacagccacgatgatcgcgggctccct 294
         Sbjct: 60 agccacagccacgatgatcgcgggctccct 89
\square >gi|32880108|gb|BT009883.1| Synthetic construct Homo sapiens tumor necrosis fact
         superfamily, member 21 mRNA, partial cds
        Length = 1968
 Score = 144 bits (72), Expect = 1e-31
 Identities = 87/90 (96%), Gaps = 2/90 (2%)
 Strand = Plus / Plus
Query: 206 atggggacctctccgagcagcagcaccgccctcggcctcctgcm-ccgcatcgcccgccg 264
         Sbjct: 1
         atggggacctctccgagcagcagcaccgccctcg-cctcctqcagccqcatcqccqccq 59
Query: 265 agccacagccacgatgatcgcgggctccct 294
         Sbjct: 60 agccacagccacgatgatcgcgggctccct 89
[]>gi|30583678|gb|BT007420.1| GUE Homo sapiens tumor necrosis factor receptor su
         mRNA, complete cds
        Length = 1968
Score = 144 bits (72), Expect = 1e-31
Identities = 87/90 (96%), Gaps = 2/90 (2%)
Strand = Plus / Plus
Query: 206 atggggacctctccgagcagcagcaccgccctcggcctcctgcm-ccgcatcgcccqccq 264
         atggggacctctccgagcagcagcaccgccctcg-cctcctgcagccgcatcgcccgccg 59
```

```
Query: 265 agccacagccacgatgatcgcgggctccct 294
        Sbjct: 60 agccacagccacgatgatcgcgggctccct 89
☐ >gi|55626753|ref|XM 518523.1| PREDICTED: Pan troglodytes tumor necrosis factor r
        superfamily, member 21 (LOC462749), mRNA
       Length = 2364
Score = 136 bits (68), Expect = 3e-29
Identities = 86/90 (95\%), Gaps = 2/90 (2\%)
Strand = Plus / Plus
Query: 206 atggggacctctccgagcagcagcaccgccctcggcctcctgcm-ccgcatcgcccgccg 264
        atggggacctctgcgagcagcagcaccgccctcg-cctcctgcagccgcatcgcccgccg 59
Sbjct: 1
Query: 265 agccacagccacgatgatcgcgggctccct 294
        Sbjct: 60 agccacagccacgatgatcgcgggctccct 89
☐ >gi|15870715|emb|AJ326321.1|HSA326321
                               Homo sapiens genomic sequence surrounding
       Length = 713
Score = 122 \text{ bits (61)}, Expect = 5e-25
Identities = 85/93 (91%), Gaps = 1/93 (1%)
Strand = Plus / Plus
Query: 15 gcmggtgctgagcgcccctagngcctcccttgccgcctccctctctctcccggccgtagc 74
        Query: 75 agtgcacat-gngtgttggaggtagatgggctc 106
       Sbjct: 77 aatgcacatggggtgttggaggtagattggctc 109
Length = 2239
Score = 96.3 bits (48), Expect = 4e-17
Identities = 87/97 (89%), Gaps = 3/97 (3%)
Strand = Plus / Plus
Query: 73 gcagtgcacatgng-tgttggaggtagatgggctc-ccggccgggaggcggcggtggatg 130
        Sbjct: 129 gcagtgcacatgggctgctggaggtagatgggctcaccgcccgtqagqcqqcqqtqqatq 188
```

```
Query: 131 cggcgctgggcag-mgcagccgccgattccagctgcc 166
        Sbjct: 189 cggcgctgggcagaaacagccaccgattccagctgcc 225
Score = 56.4 bits (28), Expect = 4e-05
Identities = 91/110 (82%), Gaps = 2/110 (1%)
Strand = Plus / Plus
Query: 186 tgcgagtccccggttcagccatggggacctctccgagcagcagcaccgcctcggcctcc 245
        Sbjct: 252 tgcgagccccgggctcagccatggggacctccgcaagcagcatcaccgccctc-gcctct 310
Query: 246 tgcm-ccgcatcgccgcgagccacagccacgatgatcgcgggctccct 294
        Sbjct: 311 tgcagccgcatcgccggccaagttggagccacgatggtcgccggctccct 360
                         G U Mus musculus tumor necrosis factor receptor su
☐ >qi|31341673|ref|NM 178589.2|
        (Tnfrsf21), mRNA
       Length = 3628
Score = 88.3 bits (44), Expect = 9e-15
Identities = 86/97 (88%), Gaps = 3/97 (3%)
Strand = Plus / Plus
Query: 73 gcagtgcacatgng-tgttggaggtagatgggctc-ccggccgggaggcggcggtggatg 130
        Sbjct: 302 gcaqtgcacatgggctgctggaggtagatgggctcaccgcccgtgaggcggcggtggatg 361
Query: 131 cggcgctgggcag-mgcagccgccgattccagctgcc 166
        Sbjct: 362 cggcgctgggcagaaacagccaccaattccagctgcc 398
enriched library, clone:A830037A05 product:Death
        Receptor 6, full insert sequence
       Length = 3628
Score = 88.3 bits (44), Expect = 9e-15
Identities = 86/97 (88%), Gaps = 3/97 (3%)
Strand = Plus / Plus
Query: 73 gcagtgcacatgng-tgttggaggtagatgggctc-ccggccgggaggcggcggtggatg 130
        Sbjct: 302 gcagtgcacatgggctgctggaggtagatgggctcaccgcccgtgaggcggcggtggatg 361
Query: 131 cggcgctgggcag-mgcagccgccgattccagctgcc 166
```

Sbjct: 362 cggcgctgggcagaaacagccaccaattccagctgcc 398

```
GUE Mus musculus tumor necrosis factor receptor su
\square > qi | 16741136 | qb | BC016420.1 |
        mRNA (cDNA clone MGC:25901 IMAGE:4220624), complete cds
       Length = 2748
 Score = 86.3 bits (43), Expect = 4e-14
 Identities = 85/96 (88%), Gaps = 3/96 (3%)
Strand = Plus / Plus
Query: 73 gcagtgcacatgng-tgttggaggtagatgggctc-ccggccgggaggcggcggtggatg 130
        Sbjct: 249 gcagtgcacatgggctgctggaggtagatgggctcaccgcccgtgaggcggcggtggatg 308
Query: 131 cggcgctgggcag-mgcagccgccgattccagctgc 165
        Sbjct: 309 cggcgctgggcagaaacagccaccaattccagctgc 344
library, clone:9030604G24 product:Death Receptor 6, full
        insert sequence
       Length = 3597
Score = 82.3 bits (41), Expect = 6e-13
Identities = 71/79 (89%), Gaps = 2/79 (2%)
Strand = Plus / Plus
Query: 90 tggaggtagatgggctc-ccggccgggaggcggtggatgcggcgctgggcag-mgca 147
        Sbjct: 295 tggaggtagatgggctcaccgcccgtgaggcggcggtggatgcggcgctgggcagaaaca 354
Query: 148 gccgccgattccagctgcc 166
        Sbjct: 355 gccaccaattccagctgcc 373
☐ >qi|15877202|emb|AJ332784.1|HSA332784
                                Homo sapiens genomic sequence surrounding
       NL6-DB13C
       Length = 670
Score = 52.4 bits (26), Expect = 6e-04
Identities = 47/55 (85%)
Strand = Plus / Plus
Query: 15 gcmggtgctgagcgcccctagngcctcccttgccgcctccctctctgcccggcc 69
```

```
□ >qi|17432071|emb|AL646086.1| □ Ralstonia solanacearum GMI1000 megaplasmid, compl
            11/11
         Length = 92509
Score = 44.4 bits (22), Expect = 0.15
Identities = 22/22 (100%)
Strand = Plus / Plus
Query: 222
           gcagcagcaccgcctcggcct 243
           Sbjct: 11349 gcagcagcaccgcctcggcct 11370
                            D Idiomarina loihiensis L2TR, complete genome
☐ >gi|56178122|gb|AE017340.1|
         Length = 2839318
Score = 42.4 bits (21), Expect = 0.59
Identities = 21/21 (100%)
Strand = Plus / Plus
Query: 107
           ccggccgggaggcggcggtgg 127
            Sbjct: 25553 ccggccgggaggcggtgg 25573
□ >gi|22795236|gb|AC127460.2| □ Homo sapiens chromosome 5 clone RP11-689B23, compl
        Length = 115293
Score = 42.4 bits (21), Expect = 0.59
Identities = 24/25 (96%)
Strand = Plus / Plus
Query: 100
           tgggctcccggccgggaggcggcgg 124
           Sbjct: 37541 tgggctcccggccggcaggcggcgg 37565
Sqi|14625084|gb|AC084380.8|AC084380 DOTYZA SATIVA CHROMOSOME 3 BAC OSJNBa0090P2
           sequence
         Length = 126164
Score = 42.4 bits (21), Expect = 0.59
Identities = 21/21 (100%)
Strand = Plus / Minus
Query: 46
           gccgcctccctcctctgcccg 66
           Sbjct: 35828 gccgcctccctcctctgcccg 35808
```

```
□>qi|33879154|qb|BC021165.2| GU Homo sapiens zinc finger protein 503, mRNA (cDNA
          IMAGE: 3830632), complete cds
         Length = 2446
 Score = 40.4 bits (20), Expect = 2.3
 Identities = 23/24 (95%)
 Strand = Plus / Plus
Query: 110 gccgggaggcggcggtggatgcgg 133
         Sbjct: 670 gccgggaggcggcggtggaggcgg 693
□>gi|33870187|gb|BC013011.2| GU Homo sapiens zinc finger protein 503, mRNA (cDNA
          IMAGE: 3604473), complete cds
         Length = 2446
 Score = 40.4 bits (20), Expect = 2.3
 Identities = 23/24 (95%)
 Strand = Plus / Plus
Query: 110 gccgggaggcggcggtggatgcgg 133
          Sbjct: 670 gccgggaggcggcggtggaggcgg 693
scleroderma antigen; RNA, U3 small nucleolar
         interacting protein 1 (LOC363563), mRNA
         Length = 882
 Score = 40.4 bits (20), Expect = 2.3
 Identities = 20/20 (100%)
 Strand = Plus / Minus
Query: 37 gcctcccttgccgcctccct 56
        Sbjct: 66 gcctcccttgccgcctccct 47
\square > gi|25453488|gb|AY057439.2| Azotobacter vinelandii GroEL (groEL) and group II in
          open reading frame genes, complete cds; and probable
          pseudouridylate synthase (rsuA) gene, partial cds
        Length = 5054
Score = 40.4 bits (20), Expect = 2.3
Identities = 20/20 (100%)
Strand = Plus / Minus
Query: 170 cgcgccccggccaccttgcg 189
          1111111111111111111
Sbjct: 2001 cgcgcccggccaccttgcg 1982
```

```
☐ >qi|52421840|qb|CP000011.1| ☐ Burkholderia mallei ATCC 23344 chromosome 2, compl
         Length = 2325379
 Score = 40.4 bits (20), Expect = 2.3
 Identities = 20/20 (100%)
 Strand = Plus / Plus
Query: 122
             cggtggatgcggcgctgggc 141
             Sbjct: 943204 cggtggatgcggcgctgggc 943223
 Score = 40.4 bits (20), Expect = 2.3
 Identities = 20/20 (100%)
 Strand = Plus / Plus
Query: 122
             cggtggatgcggcgctgggc 141
             Sbjct: 468147 cggtggatgcggcgctgggc 468166
 Score = 34.4 bits (17), Expect =
 Identities = 17/17 (100%)
 Strand = Plus / Minus
Query: 218
              ccgagcagcagcaccgc 234
              111111111111111111
Sbjct: 1853112 ccgagcagcagcaccgc 1853096
Score = 34.4 bits (17), Expect =
                                  149
Identities = 17/17 (100%)
Strand = Plus / Minus
Query: 218
              ccgagcagcagcaccgc 234
              1111111111111111
Sbjct: 1526326 ccgagcagcagcaccgc 1526310
Score = 32.4 bits (16), Expect =
                                  594
Identities = 16/16 (100%)
Strand = Plus / Plus
Query: 218
              ccgagcagcagcaccg 233
```

Sbjct: 2150293 ccgagcagcagcaccg 2150308